Concentrative Transport of Antifolates Mediated by the Proton-Coupled Folate Transporter (SLC46A1); Augmentation by a HEPES Buffer

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ABSTRACT

The proton-coupled folate transporter (PCFT) is ubiquitously expressed in solid tumors to which it delivers antifolates, particularly pemetrexed, into cancer cells. Studies of PCFT-mediated transport, to date, have focused exclusively on the influx of folates and antifolates. This article addresses the impact of PCFT on concentrative transport, critical to the formation of the active polyglutamate congeners, and at pH levels relevant to the tumor microenvironment. An HeLa-derived cell line was employed, in which folate-specific transport was mediated exclusively by PCFT. At pH 7.0, there was a substantial chemical gradient for methotrexate that decreased as the extracellular pH was increased. A chemical gradient was still detected at pH 7.4 in the usual HEPES-based transport buffer in contrast to what was observed in a bicarbonate/CO₂-buffered medium. This antifolate gradient correlated with an alkaline intracellular pH in the former (pH 7.85), but not the latter (pH 7.39), buffer and was abolished by the protonophore carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone. The gradient in HEPES buffer at pH 7.4 was the result of the activity of Na⁺/H⁺ exchanger(s); it was eliminated by inhibitors of Na⁺/H⁺ ATPase. An antifolate chemical gradient was also detected in bicarbonate buffer at pH 6.9 versus 7.4, also suppressed by carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone. When the membrane potential is considered, PCFT generates substantial transmembrane electrochemical-potential gradients at extracellular pH levels relevant to the tumor microenvironment. The augmentation of intracellular pH, when cells are in a HEPES buffer, should be taken into consideration in studies that encompass all proton-coupled transporter families.

Introduction

The proton-coupled folate transporter (PCFT) is widely expressed in human epithelial cancers (Zhao et al., 2004a; Desmoulin et al., 2011). Its low optimum pH favors a role in the delivery of antifolates within the hypoxic, acidic microenvironment of tumors (Qiu et al., 2006; Zhao and Goldman, 2013b). This may be particularly relevant to pemetrexed, an agent that has a high affinity for this transporter, and may be the reason why tumor cells that develop resistance to methotrexate (MTX) owing to loss of reduced folate carrier function, retain sensitivity to pemetrexed (Zhao et al., 2004b; Chattopadhyay et al., 2006). Low expression of PCFT strongly correlates with poor outcome in patients with mesothelioma treated with pemetrexed (Giovannetti et al., 2017). PCFT is the sole carrier-mediated route of transport of a new generation of folate analogs that are inhibitors of purine synthesis (Matherly et al., 2018). PCFT’s major physiologic role is in the intestinal absorption of folates and folate transport across the choroid plexus into the cerebrospinal fluid (Qiu et al., 2006; Zhao et al., 2009; Visentin et al., 2014). Mutations in the PCFT gene that result in loss of function of the protein are the molecular basis for the rare autosomal recessive disorder, hereditary folate malabsorption (OMIM229050) in which both functions are impaired (Qiu et al., 2006; Kronn and Goldman, 2017; Zhao et al., 2017).

The major parameter of studies on PCFT-mediated transport has been the rate of transport into cells (influx). However, once in the cell MTX and pemetrexed form polyglutamate derivatives, mediated by folylpolyglutamate synthetase (FPGS), which are retained and build to high intracellular levels (Habeck et al., 1995; Zhao et al., 2004b). In the case of pemetrexed, the polyglutamate congeners are synthesized much more rapidly than MTX by FPGS and, in contrast to the monoglutamate, are the active derivatives that inhibit tetrahydrofolate cofactor-dependent enzymes required for de novo purine and thymidylate synthesis (Shih et al., 1997; Chattopadhyay et al., 2007). The rate and extent of polyglutamate formation is dependent upon the concentration of free drug achieved within the intracellular compartment. Hence, understanding the properties of PCFT-mediated transport that govern the transmembrane antifolate gradient and the level of free drug achieved in the intracellular compartment is critical from the pharmacological perspective. This is of
particular importance under the acidic conditions that exist within the microenvironment of tumors.

The present study focuses on the characterization of net transport of MTX, in particular, concentrative transport as a function of the pH gradient across the cell membrane. In the course of these studies chemical gradients for MTX were observed at neutral extracellular pH in Hela cells studied in a HEPES buffer system usually employed for transport studies. To understand the basis for this transport, and to exploit this phenomenon to better characterize PCFT-mediated net transport, studies were undertaken with an HeLa cell line transfected to express a high level of PCFT so that this transporter would be the dominant route of antifolate transport across the cell membrane. The data characterize PCFT-mediated concentrative transport for both MTX and pemetrexed generated by an inward proton gradient at modest extracellular acidic pH levels or under conditions of mild intracellular alkalinization.

**Materials and Methods**

**Key Chemicals.** [3,5,7-3H(N)]MTX and generally tritium-labeled pemetrexed was obtained from Moravek Inc. (Brea, CA). Carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazide (FCCP), 5-(N-ethyl-N-isopropyl)amiloride (EIPA), monensin, ouabain, and nigericin were purchased from Sigma-Aldrich (St. Louis, MO). BCCF-AM was obtained from Thermo Fisher Scientific (Waltham, MA).

**Cells, Culture Conditions, and Transfections.** R1-11-PCFT-h cells are stable PCFT transfectants that originate from HeLa cells and express high levels of PCFT but lack constitutive expression of reduced folate carrier (RFC) (Zhao et al., 2013). R1-11-RFC-6 cells are stable transfectants that also originate from HeLa cells and express constitutive levels of RFC but lack expression of PCFT (Zhao et al., 2008). These cells were derived from clonal lines and this, along with the very well defined antifolate IC50s, is consistent with homogeneous expression of PCFT or RFC (Zhao et al., 2008, 2014). R1-11 cells that do not express RFC and PCFT (Zhao et al., 2004a; Diop-Bove et al., 2009) were recipients of both stable transfections. All these cells were grown in RPMI-1640 medium supplemented with 10% fetal bovine serum, 100 IU/ml of penicillin, and 100 μg/ml streptomycin. Hygromycin (0.3 mg/ml) was added to the growth medium for R1-11-RFC-6 cells; zeocin (0.1 mg/ml) was included in the growth medium for R1-11-RFC-6 cells to maintain carrier expression. For transient transfections, RFC or PCFT cDNAs constructed in pcDNA 3.1 were used as the expression vectors. R1-11 cells were seeded in 20 ml Low Background Lipofectamine 2000 (Invitrogen, Carlsbad, CA). Assessment of net MTX uptake was performed 2 days after transfection.

**Membrane Transport.** Two different buffer systems, previously employed in this laboratory, were used for transport studies: 1) HEPES-buffered saline (HBS; 20 mM HEPES, 5 mM dextrose, 140 mM NaCl, 5 mM KCl, 2 mM MgCl2) adjusted to pH 7.0, 7.2, 7.4, 7.6, 7.8, or 8.0 and 2) folate-free serum-free RPMI-1640 medium containing 24 mM sodium bicarbonate at pH 7.3. The pH of the RPMI medium was adjusted to 6.9 by reducing the sodium bicarbonate concentration to 7.2 mM (Zhao et al., 2004a). Serum-free RPMI medium is used in these studies to exclude binding of tritiated antifolates to serum proteins. Cells in both RPMI preparations were maintained in a 37°C incubator under an atmosphere of 5% CO₂ for at least 4 hours before experiments were initiated. For measurements of net uptake, cells that had been seeded in the glass scintillation vials and reached early confluency were washed twice with HBS (pH 7.4) or RPMI and incubated for 20 minutes either in HBS in a 37°C water bath or in RPMI at 37°C in a CO₂ incubator. Uptake was initiated by replacing the buffer with 0.5 ml prewarmed HBS or RPMI containing the desired concentration of tritiated drug. For assessment of the effects of FCCP, EIPA, ouabain, or monensin on net uptake, a small volume of 1000× stock solution either in dimethyl sulfoxide or ethanol was diluted in transport buffer with tritiated drug. Uptake was continued for 40 minutes at 37°C, unless otherwise specified, following which 5 ml of ice-cold HBS (pH 7.4) was added, the cells washed three times, then digested in 0.5 ml of 0.2 M NaOH at 65°C for 1 hour. Radioactivity in 0.4 ml of lysate was determined on a liquid scintillation spectrometer and normalized to the protein level obtained with the BCA Protein Assay (Pierce Protein Biology/Thermo Fisher Scientific). Net uptake of antifolates was expressed as picomoles per milligram protein or percentages of control. For some experiments units of picomoles per milligram protein were converted to intracellular concentration (micromolar) on the basis of the conversion factor 8.73 pmol/mg protein per micromolar (Sharif et al., 1998).

**Results**

**Transmembrane MTX Gradients as a Function of Extracellular pH.** Studies were undertaken to assess the extent to which there is concentrative MTX transport as the pH of HEPES-based saline (HBS) increases from mildly acidic to the basic pH range. An HeLa cell line, R1-11-PCFT-h, was employed that lacks RFC expression but stably expresses a high level of PCFT. For these studies chemical gradients for MTX were generated by an inward proton gradient at modest extracellular acidic pH levels or under conditions of mild intracellular alkalinization.

**Intracellular pH Measurements.** These measurements were obtained as previously described (Zhao et al., 2013). Briefly, R1-11-PCFT-h cells grown in glass-bottom dishes (MatTek, Ashland, MA) were loaded with BCECF-AM, a pH indicator, (10 μM, Molecular Probes/Invitrogen; Thermo Fisher Scientific) in culture media for 45 minutes at 37°C in a humidified 5% CO₂ incubator. Cells were then rinsed with prewarmed HBS or 5% CO₂-saturated RPMI and imaged at 37°C with an Orea-EM camera (Hamamatsu, Japan) attached to a Nikon TE2000 microscope. Changes in BCECF fluorescence intensity emitted at two excitation wavelengths (480 and 450 nm) were acquired at 1.0 Hz using filters and shutter (Lambda DG-4; Sutter Instruments, Novato, CA) driven by a computer through MetaFluor software (Molecular Devices, Silicon Valley, CA). Intracellular pH was calibrated for each dish with MOPS-buffered pH solutions of 140 mM potassium chloride and 20 μM nigericin at pH 8.0 or pH 7.0.

**Discussion**

The results demonstrate that concentrative MTX transport is a general phenomenon to better characterize PCFT-mediated net trans-
Fig. 1. The MTX transmembrane gradient as a function of extracellular pH in cells incubated in HBS. After a 20-minute preincubation of R1-11-PCFT-h cells in HBS (pH 7.40), [3H]MTX was added to achieve a concentration of 100 μM, and uptake continued for 40 minutes in the presence or absence of 10 μM FCCP in HBS over a range of pH values. Net uptake of MTX in HBS (pH 7.40) in the absence of FCCP was set as 100% as a basis for comparison. The percentage of uptake in the presence vs. absence of FCCP is indicated above the bars for each pH. The intracellular MTX concentrations in units of micromolarity are indicated in the right Y-axis. The horizontal line indicates the extracellular MTX concentration at 100 μM. Data are the mean ± S.D. from three separate experiments.

Analysis of Net PCFT-Mediated MTX Transport When the Extracellular pH Is Neutral; Impact of the Buffer. To assess whether HEPES-buffered HBS might impact upon PCFT-mediated transmembrane MTX gradients, net transport was compared 10–50 minutes after addition of [3H]MTX in two buffer systems: 1) HBS (pH 7.40), which is HEPES-buffered and does not contain NaHCO3, and 2) folate- and serum-free RPMI 1640 medium buffered with 24 mM bicarbonate and maintained in an atmosphere of 5% CO2 at an extracellular pH of ~7.3. As indicated in Fig. 2, net MTX uptake reached 75% of steady state in HBS by 25 minutes, rising to a steady state by 40 minutes, whereas MTX uptake in RPMI was at steady state by 25 minutes at one-third the level in HBS. Inclusion of 10 μM FCCP markedly decreased net MTX in cells in HBS. FCCP resulted in a smaller decrease in net MTX in the RPMI buffer. Hence, the difference in net MTX levels in cells maintained in the different buffers was substantially decreased when the proton gradient was abolished by FCCP.

When the intracellular MTX level is expressed in micro-moles per liter, as in the right Y-axis of Fig. 2, it can be seen there was an approximately 3.1 chemical gradient for MTX in HBS that disappeared in the presence of FCCP. In RPMI there was no chemical gradient in the absence of FCCP and the net level was about half of that of the extracellular concentration in the presence of FCCP. Hence, net MTX transport was concentrically primary in HBS at pH 7.4. This difference in net MTX transport correlated with the difference in intracellular pH as assessed with BCECF dye. As indicated in Fig. 3, the intracellular pH in HBS (7.85 ± 0.18 (mean ± S.D.), n = 6, Fig. 3A) was ~0.46 pH units higher than cells in RPMI [pH 7.39 ± 0.19 (mean ± S.D.), n = 3, Fig. 3B (P = 0.010, unpaired two-tailed t test)]. Hence, despite the fact that the extracellular pH was nearly the same in the two buffers, there was a substantial inward gradient in HBS, whereas a proton gradient could not be detected in cells in RPMI.

Mechanism of Intracellular Alkalization in HEPES-Buffered Medium. Studies were undertaken to determine the basis for the increased intracellular pH detected in the HBS buffer. The initial focus was on Na+/H+ antiporters that contribute to the maintenance of intracellular pH by exporting protons in exchange for sodium ions. EIPA (10 μM), a specific inhibitor of Na+/H+ antiporters (Masereel et al., 2003), decreased the net MTX level in cells in HBS by 77%, whereas EIPA had no effect on the net MTX level in cells in RPMI (Fig. 4A). Likewise, a decrease in the Na+/K+ gradients by inhibition of the Na+/K+ ATPase with 10 μM ouabain also markedly reduced net MTX uptake in cells in HBS. Finally, although monensin (50 μM), a sodium/proton exchanger (Nakazato and Hatano, 1991), did not alter net MTX uptake in HBS, the net MTX level more than doubled in RPMI. These observations suggested that the inward proton gradient in cells in HBS was generated by Na+/H+ antiporters and sustained by the ATPase-dependent sodium gradient.

To explore the basis for the augmentation of net MTX uptake by monensin in cells in the RPMI buffer, the impact of the addition of other agents in the presence of monensin was assessed. As indicated in Fig. 4B, net MTX uptake in HBS was ~2.3-fold higher than in RPMI in these experiments. Monensin at 50 μM increased net MTX uptake in RPMI to the level achieved in HBS. This was unchanged in the presence of EIPA but was abolished by FCCP or ouabain declining to the level with FCCP alone. In contrast to the marked reduction in MTX uptake induced by EIPA in the absence of monensin cells in HBS, EIPA had no effect at all on MTX uptake in the presence of monensin either in cells in HBS or RPMI.

Potential Role of ABC Exporters in the Modulation of Net MTX Levels. The net level of free intracellular folates and antifolates is determined by the balance between the carriers PCFT or RFC, which generates an inward gradient, and by several ABC exporters, such as multidrug-resistant proteins that pump folates/antifolates out of cells (Assaraf, 2006). Hence, changes in net MTX levels observed with these inhibitors might be explained, in part, by their effects on ABC exporters. The high level of expression of PCFT in these cells was designed to decrease the impact of ABC exporters. However, further studies were designed to assess possible changes in ABC exporters that impact upon the assessment of PCFT-mediated net antifolate levels. Effects of alterations in ABC exporter function should also be observed under...
conditions in which the net level of intracellular antifolates is mediated solely by RFC, a folate/organic phosphate antiporter (Assaraf, 2006). To evaluate this, the impact of these agents was assessed in R1-11-RFC-6 cells that do not express PCFT but stably express RFC at levels comparable to that of wild-type HeLa cells. Any contribution of the ABC exporters to the net MTX level should overwhelm the low level of MTX cycling mediated by RFC under these conditions. As indicated in Fig. 5A, there was no significant difference in RFC-mediated steady-state MTX levels in RPMI and, beyond a very small decrease (15%, \( P = 0.029 \), paired two-tailed \( t \) test) in the EIPA-treated cells, none of these agents altered net MTX uptake mediated by RFC in the cells in HBS. Hence, the data were consistent with effects of these agents on R1-11-PCFT-h cells owing specifically to effects on PCFT-mediated transport and were unrelated to alterations in ABC exporter activities.

To further exclude the possibility that the effects of FCCP, EIPA, or ouabain on net MTX uptake observed in PCFT and RFC stable transfectants were related to possible differences in the expression of ABC exporters, the impact of these inhibitors on net MTX uptake was assessed in RFC- and PCFT-transient transfectants using the same promoter and exactly the same conditions. Under these conditions, expression of ABC exporters in both transient transfectants should have been comparable. As indicated in Fig. 5B, transfections of either RFC or PCFT increased net MTX uptake compared with the mock transfection. However, FCCP, EIPA, and ouabain did not significantly alter net MTX uptake in RFC transient transfects, whereas these agents reduced MTX net uptake by more than 50% in PCFT transient transfectants (\( P < 0.05 \)). The lesser effects on net MTX uptake by these reagents observed in PCFT-transient transfectants compared with the PCFT-stable transfectants is
probably related to the difference in MTX net uptake (∼300 pmol/mg vs. ∼3000 pmol/mg protein for transient and stable transfectants, respectively). As observed with the stable transfectants, monensin had no effect on net MTX uptake for either RFC or PCFT transient transfectants. The lack of an impact by these inhibitors on net MTX uptake mediated by RFC in the transient transfectants is additional evidence that the effects of FCCP, EIPA, and ouabain on net MTX uptake are entirely the result of changes in PCFT-mediated transport and are not related to alterations in expression of ABC exporters.

Net Transport of Pemetrexed, Effects of Buffers Monensin and FCCP. Net transport of pemetrexed was also assessed in HBS or RPMI in the presence or absence of monensin and FCCP. Pemetrexed forms polyglutamate derivatives that are retained within cells and accumulate much more rapidly than MTX. This complicates discrimination between monoglutamate, free within the intracellular water, and tritiated polyglutamylated congeners that cannot exit the cells. To address this, in preparation for experiments, R1-11-PCFT-h cells were grown with 0.1 mM folic acid overnight to allow accumulation of reduced folate derivatives that are good substrates for FPGS and compete with, and suppress, the polyglutamation of pemetrexed. Net [3H]pemetrexed uptake was then assessed at 50 μM (the influx Ki for pemetrexed at pH 7.4 is 12–15 μM) for 40 minutes. This concentration is also relevant to pemetrexed blood levels after intravenous administration of the drug (Latz et al., 2006). As indicated in Fig. 6, and as was observed for MTX, net pemetrexed uptake in HBS was 2.5-fold greater than in RPMI at pH 7.4. Monensin increased pemetrexed net transport in RPMI by a factor of about 2.5 but did not affect net pemetrexed uptake in HBS. FCCP decreased the net pemetrexed level to ∼25 μM, one-half the extracellular level in either buffer, and also suppressed the increase observed with monensin in RPMI.

Analysis of Net Intracellular MTX Transport in RPMI at pH 7.3 versus 6.9. To better simulate conditions within the microenvironment of solid tumors in vivo, using a physiologic buffered system, MTX levels were evaluated in cells in RPMI at pH 7.3 and RPMI adjusted to a pH of 6.9 by decreasing the bicarbonate concentration to 7.2 mM in a 5% CO2 atmosphere, in the presence or absence of FCCP, ouabain, EIPA, or monensin. As indicated in Fig. 7, the net intracellular MTX level nearly doubled at pH 6.9 versus 7.3. When the membrane potential is considered, this represents a substantial electrochemical-potential gradient for MTX. FCCP resulted in a decline in the net intracellular MTX levels to a comparable level in both buffers at approximately half that at
pH 7.3. The addition of monensin resulted in a 2.2-fold increase in the MTX level at both pH 6.9 and 7.3 that was abolished by FCCP. As a result, a combination of lower extracellular pH and the presence of monensin resulted in a 2.2-fold augmentation in the intracellular MTX level in RPMI. Extracellular pH and the presence of monensin resulted in a 2-fold augmentation in the intracellular MTX level in RPMI (pH 7.3) that was abolished by FCCP. As a result, a combination of lower extracellular pH and the presence of monensin resulted in a 2-fold augmentation in the intracellular MTX level in RPMI. Extracellular pH and the presence of monensin resulted in a 2-fold augmentation in the intracellular MTX level in RPMI (pH 7.3) is indicated as 100%. Data are the mean ± S.D. from three independent experiments.

**Discussion**

Studies of solute transporters usually focus on initial uptake properties. This may provide sufficient information for equilibrating systems that do not generate transmembrane gradients. However, the properties of transporters that generate gradients by the coupled flow of other ions or organic solutes is more complex. The free intracellular substrate level achieved may not be critical for some pharmacological agents, particularly when influx is rate-limiting to an important intracellular event. However, this parameter of the drug-cell interaction controlled by the membrane transporter is considerably more important when substantial levels of free intracellular drug are necessary to drive a subsequent biochemical step critical for drug retention, metabolism to an active derivative(s), or, in the case of a physiologic substrate, its normal cellular metabolism. This is the case for virtually all antifolate drugs now in clinical use, in particular pemetrexed, which requires polyglutamation for its activity (Habeck et al., 1995; Shih et al., 1997), and a new generation of glycaminide ribonucleotide transformylase inhibitors with an equally high affinity for PCFT but a very low affinity for RFC (Desmoulin et al., 2012; Hou et al., 2017). The latter agents are only active in their polyglutamate forms.

Studies of the concentrative potential of PCFT, in cells that typically exist in the normal systemic environment where the pH is maintained at 7.4, are complicated by the rapid acidification of many cell types after exposure to acidic conditions. However, we demonstrate here that modest alkalization of cells, under conditions in which the extracellular pH is at the normal systemic level, as occurs in HEPES-buffered media, results in a chemical gradient for MTX and pemetrexed. Hence, although the intracellular pH is essentially the same as the extracellular pH in RPMI medium, the intracellular pH is 0.46 pH units higher when cells are incubated in a HEPES buffer under conditions used for most transport studies. The chemical gradients for MTX and pemetrexed are abolished with FCCP, suggesting that in the HEPES buffer they are driven, entirely, by a pH gradient across the cell membrane.

Hela cells have a substantial negative membrane potential, ~50 mV, and MTX and pemetrexed carry negative charges at each carboxyl group in the glutamate moiety. Hence, at thermodynamic equilibrium, when there is no electrochemical-potential difference across the cell membrane, the concentration of these antifolates within the cell should be far below their extracellular concentration. For instance, assuming MTX to be a bivalent anion and the membrane potential to be ~50 mV (Stein et al., 1996), a reverse gradient is expected of ~0.3 [MTX]/[MTX]o. Accordingly, when the ratio of intracellular-to-extracellular concentration is one, there is in fact a 3-fold electrochemical-potential gradient across the cell membrane. This is consistent with the observation that the addition of FCCP results in intracellular MTX levels that are below the extracellular concentration. It is also of interest that FCCP appears to decrease net intracellular MTX in the bicarbonate-buffered system even under conditions in which there should not be a proton gradient. Beyond abolishing the proton gradient across the cell and mitochondrial membranes, FCCP also lowers the membrane potential (Park et al., 2002) which should increase, rather than decrease, net MTX levels. As FCCP also inhibits oxidation, this too would increase cellular MTX levels by decreasing ATP-dependent multidrug-resistant protein activity. Finally, although there are solute transporters that exhibit proton coupling at acidic pH and Na+ coupling at neutral pH (Smith et al., 2005; Young et al., 2013), so that transmembrane gradients are present at neutral pH, there is no evidence that PCFT is Na+/H+ coupled at neutral pH and thus not responsible in part for the gradients observed (Zhao et al., 2013).

The data indicate that the chemical gradients observed for MTX and pemetrexed in HBS buffer result from a pH gradient generated by Na+/H+ antiporter(s). Maintenance of the Na+ gradient, which drives the Na+/H+ exchange, requires Na+/K+ ATPase activity (Fig. 8). Hence, both EIPA and ouabain completely eliminate the antifolate gradient. Interestingly, monensin, an ionophore that functions as a Na+/H+ exchanger (Nakazato and Hatano, 1991), does not further increase net transport of MTX in cells in HBS, presumably because the proton gradient already present was generated by the constitutive Na+/H+ exchanger(s). On the other hand, the monensin-induced MTX gradient in cells in RPMI is consistent with monensin acting as a Na+/H+ exchanger to result in
a proton gradient that is abolished by FCCP and ouabain but is unaffected by EIPA. Although it would have been of interest to determine whether these changes in net MTX uptake are associated with changes in the antitumor activities of the antifolates, the intrinsic toxicity of these agents precluded an assessment of antifolate growth inhibition or cytotoxicity under these conditions.

Steady-state levels of folates and antifolates are determined by a variety of processes. Under physiologic conditions at pH 7.4, RFC transports folates into and out of virtually all systemic cells and has concentrative capability through an exchange with intracellular organic phosphates, in particular, adenine nucleotides (Zhao and Goldman, 2013a). The concentrative impact of both PCFT and RFC is opposed by several ABC exporters that transport the monoglutamate (and in some cases the lower polyglutamate derivatives) of folates and antifolates out of cells (Assaraf, 2006). The experimental design in the present study precluded an RFC contribution to the net antifolate level in these cells since they express only high levels of PCFT. A potential role for the ABC exporters as a route of delivery of contemporary antifolates into cancer cells, in particular, pemetrexed, an agent with high affinity for the polyglutamation of this agent (Zhao et al., 2004b; Chattopadhyay et al., 2006). This is probably the basis for genetic distinction in the present study.

As yet incompletely understood is the role that PCFT plays as a route of delivery of contemporary antifolates into cancer cells, in particular, pemetrexed, an agent with high affinity for this transporter. It is clear that PCFT can sustain the activity of pemetrexed when RFC is deleted, complemented by the concurrent contraction of cellular folates that enhances the polyglutamation of this agent (Zhao et al., 2004b; Chattopadhyay et al., 2006). This is probably the basis for the observation that it is very difficult, if not impossible, for cells to develop resistance to this drug because of a failure of transport into cells, which would require the loss of two genetically distinct transporters. PCFT is the sole solute transporter that mediates the activity of six-substituted pyrrolo[2,3-d]pyrimidine antifolate purine synthesis inhibitors that are very poor substrates for RFC (Desmoulion et al., 2012; Hou et al., 2017). The latter is a strategy for

![Diagram](Image137x576to448x729)
circuiting the toxicity of classic antifolates that is mediated primarily by their delivery into susceptible cells of the gastrointestinal tract and bone marrow by RFC. The pemetrexed concentration in plasma after a standard intravenous dose is in the range of ∼200 µM with an initial half-life of ∼1 hour and concentrations in excess of 20 µM sustained for ∼3 hours (Latz et al., 2006). Assuming even a neutral intratumoral pH, and a pemetrexed influx Kf of 12–15 µM at pH 7.4, there should be substantial delivery and free intracellular drug levels mediated by PCFT over this interval. However, since the usual interstitial pH of tumors is acidic, PCFT-mediated pemetrexed influx and free intracellular levels would be much greater resulting in increased formation of active pemetrexed polyglutamate derivatives (Wike-Hooley et al., 1984; Tannock and Rotin, 1989; Hashim et al., 2011).

Authorship Contributions

Participated in research design: Zhao, Spray, Goldman. Conducted experiments: Zhao, Najmi, Aluri. Performed data analysis: Zhao, Najmi, Aluri, Spray, Goldman. Wrote or contributed to the writing of the manuscript: Zhao, Goldman.

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